

FORM PTO-1449

FOURTH SUPPLEMENTAL INFORMATION  
DISCLOSURE STATEMENTATTY. DOCKET NO.  
1503.1070003APPLICATION NO.  
09/934,084APPLICANTS  
Lobanov et al.FILING DATE  
August 22, 2001GROUP  
1631

## U.S. PATENT DOCUMENTS

EXAMINER INITIAL		DOCUMENT NUMBER	DATE	NAME	CLASS	SUB- CLASS	FILING DATE
	AA1						
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## FOREIGN PATENT DOCUMENTS

EXAMINER INITIAL		DOCUMENT NUMBER	DATE	COUNTRY	CLASS	SUB- CLASS	TRANSLATION
CSM	AJ1	EP 0 818 744 A2	01/1998	EPO	G06F	17/50	Yes No
	AK1	WO 93/20242	10/1993	PCT	C12Q	1/70	Yes No
	AL1	WO 95/01606	01/1995	PCT	G06F	15/42	Yes No
✓	AM1	WO 97/27559	07/1997	PCT	G06F	19/00	Yes No

## OTHER (Including Author, Title, Date, Pertinent Pages, etc.)

CSM	AN	1	Borg, Ingwer and Groenen, Patrick, <i>Modern Multidimensional Scaling Theory and Applications</i> , Springer Series in Statistics, 1997, entire book submitted.
	AO	1	Agrafiotis, D.K. et al., "Advances in diversity profiling and combinatorial series design," <i>Molecular Diversity</i> , Kluwer Academic Publishers, Vol. 4, 1999, pp. 1-22.
	AP	1	Agrafiotis, D.K. and Lobanov, V.S., "An Efficient Implementation of Distance-Based Diversity Measures Based on k-d Trees," <i>Journal of Chemical Information and Computer Science</i> , American Chemical Society, Vol. 39, No. 1, January/February 1999, pp. 51-58.
	AQ	1	Agrafiotis, D.K. and Lobanov, V.S., "Bridging The Gap Between Diversity And QSAR," <i>Abstracts of Papers Part 1: 215th ACS National Meeting</i> , American Chemical Society, March 29-April 2, 1998, p. 181-COMP.
✓	AR	1	Agrafiotis, D.K. and Jaeger, E.P., "Directed Diversity®: An Operating System For Combinatorial Chemistry," <i>Abstracts of Papers Part 1: 211th ACS National Meeting</i> , American Chemical Society, March 24-28, 1996, p. 46-COMP.

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C. M. ...

DATE CONSIDERED

March 14, 2004

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EXAMINER INITIAL		DOCUMENT NUMBER	DATE	COUNTRY	CLASS	SUB- CLASS	TRANSLATION
CSM	AJ2	WO 98/20459	05/1998	PCT	G06T	11/20	Yes No
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	AL2						Yes No
	AM2						Yes No

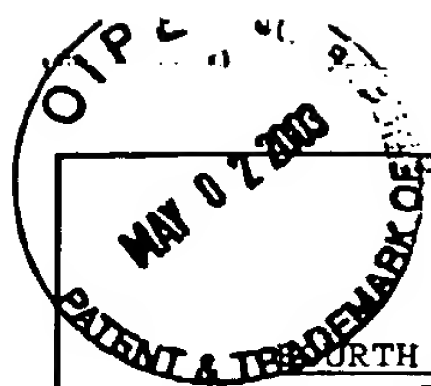
OTHER (Including Author, Title, Date, Pertinent Pages, etc.)

CSM	AN	2	Agrafiotis, D.K., "Diversity of Chemical Libraries," <i>Encyclopedia of Computational Chemistry</i> , John Wiley & Sons Ltd, Vol. 1:A-D, 1998, pp. 742-761.
	AO	2	Agrafiotis, D.K., "On the Use of Information Theory for Assessing Molecular Diversity," <i>Journal of Chemical Information and Computer Science</i> , American Chemical Society, Vol. 37, No. 3, May/June 1997, pp. 576-580.
	AP	2	Agrafiotis, D.K. et al., "Parallel QSAR," <i>Abstracts of Papers Part 1: 217th ACS National Meeting</i> , March 21-25, 1999, p. 50-COMP.
	AQ	2	Agrafiotis, D.K. et al., "PRODEN: A New Program for Calculating Integrated Projected Populations," <i>Journal of Computational Chemistry</i> , John Wiley & Sons, Inc., Vol. 11, No. 9, October 1990, pp. 1101-1110.
↓	AR	2	Agrafiotis, D.K. and Jaeger, E.P., "Stochastic Algorithms for Exploring Molecular Diversity," <i>Abstracts of Papers Part 1: 213th ACS National Meeting</i> , American Chemical Society, April 13-17, 1997, p. 16-CINF.

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	AJ3						Yes No
	AK3						Yes No
	AL3						Yes No
	AM3						Yes No

## OTHER (Including Author, Title, Date, Pertinent Pages, etc.)

CSM	AN	3	Agrafiotis, D., "Theoretical Aspects of the Complex: Arts and New Technologies," <i>Applications and Impacts Information Processing '94</i> , North-Holland, Vol. II, 1994, pp. 714-719.
	AO	3	Biswas, G. et al., "Evaluation of Projection Algorithms," <i>IEEE Transactions On Pattern Analysis And Machine Intelligence</i> , IEEE Computer Society, Vol. PAMI-3, No. 6, November 1981, pp. 701-708.
	AP	3	Bonchev, D. and Trinajstić, N., "Information theory, distance matrix, and molecular branching," <i>The Journal of Chemical Physics</i> , American Institute of Physics, Vol. 67, No. 10, November 15, 1977, pp. 4517, 4520-4533.
	AQ	3	Chang, C.L. and Lee, R.C.T., "A Heuristic Relaxation Method for Nonlinear Mapping in Cluster Analysis," <i>IEEE Transactions on Systems, Man, and Cybernetics</i> , IEEE Systems, Man, and Cybernetics Society, Vol. SMC-3, March 1973, pp. 197-200.
↓	AR	3	Cramer, R.D. et al., "Virtual Compound Libraries: A New Approach to Decision Making in Molecular Discovery Research," <i>J. Chem. Inf. Comput. Sci.</i> , American Chemical Society, Vol. 38, No. 6, November/December 1998, pp. 1010-1023.

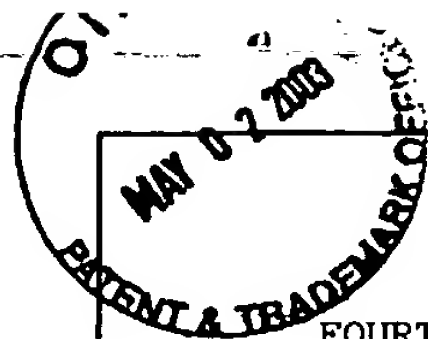
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	AJ4						Yes No
	AK4						Yes No
	AL4						Yes No
	AM4						Yes No

**OTHER (Including Author, Title, Date, Pertinent Pages, etc.)**

Cum	AN	<u>4</u>	DeMers, D. and Cottrell, G., "Non-Linear Dimensionality Reduction," <i>Advances in Neural Information Processing Systems</i> , Vol. 5, 1993, pp. 580-587.
	AO	<u>4</u>	Frey, P.W. and Slate, D.J., "Letter Recognition Using Holland-Style Adaptive Classifiers," <i>Machine Learning</i> , Kluwer Academic Publishers, Vol. 6, 1991, pp. 161-182.
	AP	<u>4</u>	Friedman, J.H., "Exploratory Projection Pursuit," <i>Journal of the American Statistical Association</i> , American Statistical Association, Vol. 82, No. 397, March 1987, pp. 249-266.
	AQ	<u>4</u>	Friedman, J.H. and Tukey, J.W., "A Projection Pursuit Algorithm for Exploratory Data Analysis," <i>IEEE Transactions on Computers</i> , IEEE Computer Society, Vol. C-23, No. 9, September 1974, pp. 881-889.
✓	AR	<u>4</u>	Garrido, L. et al., "Use of Multilayer Feedforward Neural Nets As A Display Method for Multidimensional Distributions," <i>International Journal of Neural Systems</i> , World Scientific Publishing Co. Pte. Ltd., Vol. 6, No. 3, September 1995, pp. 273-282.

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	AJ5						Yes No
	AK5						Yes No
	AL5						Yes No
	AM5						Yes No

## OTHER (Including Author, Title, Date, Pertinent Pages, etc.)

GAM	AN	5	Ghose, A.K. et al., "Prediction of Hydrophobic (Lipophilic) Properties of Small Organic Molecules Using Fragmental Methods: An Analysis of ALOGP and CLOGP Methods," <i>Journal of Physical Chemistry</i> , American Chemical Society, Vol. 102, No. 21, May 21, 1998, pp. 3762-3772.
	AO	5	Hall, L.H. and Kier, L.B., "The Molecular Connectivity Chi Indexes and Kappa Shape Indexes in Structure-Property Modeling," <i>Reviews in Computational Chemistry: Advances</i> , VCH Publishers, Inc., 1991, pp. 367-422.
	AP	5	Hecht-Nielsen, R., "Replicator Neural Networks for Universal Optimal Source Coding," <i>Science</i> , American Association for the Advancement of Science, Vol. 269, September 29, 1995, pp. 1860-1863.
	AQ	5	Hotelling, H., "Analysis of a Complex of Statistical Variables into Principal Components," <i>The Journal of Educational Psychology</i> , Warwick and York, Inc., Vol. XXIV, No. 6, September 1933, pp. 417-441.
↓	AR	5	Hotelling, H., "Analysis of a Complex of Statistical Variables into Principal Components," <i>The Journal of Educational Psychology</i> , Warwick and York, Inc., Vol. XXIV, No. 7, October 1933, pp. 498-520.

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	AJ6						No
	AK6						Yes No
	AL6						Yes No
	AM6						Yes No

**OTHER (Including Author, Title, Date, Pertinent Pages, etc.)**

C4M	AN	6	Lee, R.C.T. et al., "A Triangulation Method for the Sequential Mapping of Points from N-Space to Two-Space," <i>IEEE Transactions on Computers</i> , The Institute of Electrical and Electronics Engineers, March 1977, pp. 288-292.
	AO	6	Lipinski, C.A. et al., "Experimental and computational approaches to estimate solubility and permeability in drug discovery and development settings," <i>Advanced Drug Delivery Reviews</i> , Elsevier Science B.V., Vol. 23, 1997, pp. 3-25.
	AP	6	Lobanov, V.S. and Agrafiotis, D.K., "Intelligent Database Mining Techniques," <i>Abstracts of Papers Part 1: 215th ACS National Meeting</i> , March 29-April 2, 1998, p. 19-COMP.
	AQ	6	Lobanov, V.S. et al., "Rational Selections from Virtual Libraries," <i>Abstracts of Papers Part 1: 217th ACS National Meeting</i> , March 21-25, 1999, p. 181-COMP.
✓	AR	6	Mao, J. and Jain, A.K., "Artificial Neural Networks for Feature Extraction and Multivariate Data Projection," <i>IEEE transactions on Neural Networks</i> , IEEE Neural Networks, Vol. 6, No. 2, March 1995, pp. 296-317.

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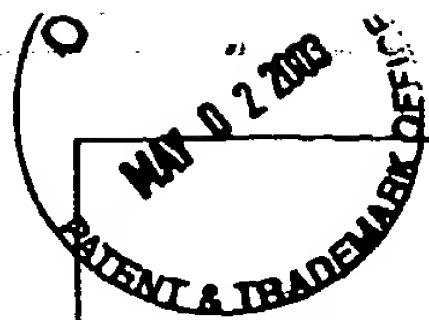
## OTHER (Including Author, Title, Date, Pertinent Pages, etc.)

GBM	AN	Z	Oja, E., "Principal Components, Minor Components, and Linear Neural Networks," <i>Neural Networks</i> , Pergamon Press Ltd., Vol. 5, 1992, pp. 927-935.
	AO	Z	Patterson, D.E. et al., "Neighborhood Behavior: A Useful Concept for Validation of 'Molecular Diversity' Descriptors," <i>Journal of Medicinal Chemistry</i> , American Chemical Society, Vol. 39, No. 16, 1996, pp. 3049-3059.
	AP	Z	Pykett, C.E., "Improving the Efficiency of Sammon's Nonlinear Mapping by Using Clustering Archetypes," <i>Electronics Letters</i> , The Institution of Electrical Engineers, Vol. 14, No. 25, December 7, 1978, pp. 799-800.
	AQ	Z	Rubner, J. and Tavan, P., "A Self-Organizing Network for Principal-Component Analysis," <i>Europhysics Letters</i> , European Physical Society, Vol. 10, No. 7, December 1, 1989, pp. 693-698.
↓	AR	Z	Sadowski, J. et al., "Assessing Similarity and Diversity of Combinatorial Libraries by Spatial Autocorrelation Functions and Neural Networks," <i>Angewandte Chemie</i> , VCH, Vol. 34, No. 23/24, January 5, 1996, pp. 2674-2677.

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	AM8						Yes No

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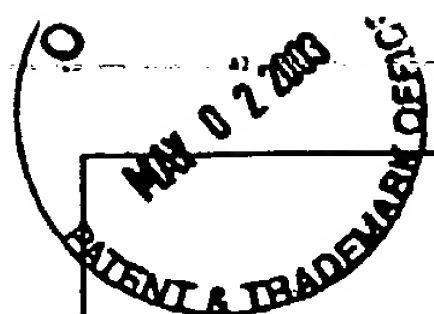
GSM ↓	AN	8	Kim, J. et al., "Multiple Neural Networks using the Reduced Input Dimension," <i>Proceedings of the International Conference on Acoustics, Speech, and Signal Processing</i> , IEEE, Vol. 2, April 19-22, 1994, pages II-601 to II-604.
	AO	8	Barnard, John M. and Downs, Geoff M., "Computer representation and manipulation of combinatorial libraries," <i>Perspectives in Drug Discovery and Design</i> , Kluwer Academic Publishers, 1997, pp. 13-30.
	AP	8	Brint, Andrew T. and Willett, Peter, "Upperbound procedures for the identification of similar three-dimensional chemical structures," <i>Journal of Computer-Aided Molecular Design</i> , ESCOM Science Publishers B.V., Vol. 2, No. 4, January 1989, pp. 311-320.
	AQ	8	Brown, Robert D. and Martin, Yvonne C., "Designing Combinatorial Library Mixtures Using a Genetic Algorithm," <i>Journal of Medicinal Chemistry</i> , American Chemical Society, Vol. 40, No. 15, 1997, pp. 2304-2313.
	AR	8	Gillet, Valerie J. et al., "The Effectiveness of Reactant Pools for Generating Structurally-Diverse Combinatorial Libraries," <i>Journal of Chemical and Information Computer Sciences</i> , American Chemical Society, Vol. 37, No. 4, 1997, pp. 731-740.

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	AJ9						Yes No
	AK9						Yes No
	AL9						Yes No
	AM9						Yes No

## OTHER (Including Author, Title, Date, Pertinent Pages, etc.)

CSM	AN	9	Gillet, Valerie J. et al., "Selecting Combinatorial Libraries to Optimize Diversity and Physical Properties," <i>Journal of Chemical Information and Computer Sciences</i> , American Chemical Society, Vol. 39, No. 1, 1999, pp. 169-177.
	AO	9	Kearsley, Simon K. et al., "Chemical Similarity Using Physiochemical Property Descriptors," <i>Journal of Chemical Information and Computer Sciences</i> , American Chemical Society, Vol. 36, No. 1, 1996, pp. 118-127.
	AP	9	Leland, Burton A. et al., "Managing the Combinatorial Explosion," <i>Journal of Chemical Information and Computer Sciences</i> , American Chemical Society, Vol. 37, No. 1, 1997, pp. 62-70.
	AQ	9	Lewis, Richard A. et al., "Similarity Measures for Rational Set Selection and Analysis of Combinatorial Libraries: The Diverse Property-Derived (DPD) Approach," <i>Journal of Chemical Information and Computer Sciences</i> , American Chemical Society, Vol. 37, No. 3, 1997, pp. 599-614.
↓	AR	9	Martin, Eric J. and Critchlow, Roger E., "Beyond Mere Diversity: Tailoring Combinatorial Libraries for Drug Discovery," <i>Journal of Combinatorial Chemistry</i> , American Chemical Society, Vol. 1, No. 1, 1999, pp. 32-45.

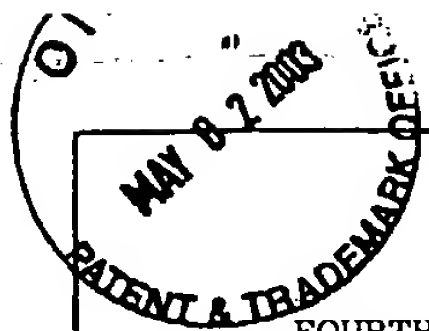
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	AK10						Yes No
	AL10						Yes No
	AM10						Yes No

## OTHER (Including Author, Title, Date, Pertinent Pages, etc.)

CAM	AN	10	Sheridan, Robert P. et al., "Chemical Similarity Using Geometric Atom Pair Descriptors," <i>Journal of Chemical Information and Computer Sciences</i> , American Chemical Society, Vol. 36, No. 1, 1996, pp. 128-136.
	AO	10	Willett, Peter et al., "Chemical Similarity Searching," <i>Journal of Chemical Information and Computer Sciences</i> , American Chemical Society, Vol. 38, No. 6, 1998, pp. 983-996.
	AP	10	Agrafiotis, Dimitris K. and Lobanov, Victor S., "Ultrafast Algorithm for Designing Focused Combinational Arrays," <i>Journal of Chemical Information and Computer Sciences</i> , American Chemical Society, 2000, Vol. 40, No. 4, pp. 1030-1038.
	AQ	10	Ajay et al., "Can We Learn To Distinguish between 'Drug-Like' and 'Nondrug-like' Molecules?" <i>J. Med. Chem.</i> , 1998, American Chemical Society, Vol. 41, No. 18, pp. 3314-3324.
↓	AR	10	Spellmeyer, D. et al., "Conformational analysis using distance geometry methods," <i>Journal of Molecular Graphics &amp; Modelling</i> , Elsevier Science, Inc., Vol. 15, No. 1, February 1997, pages 18-36.

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	AM11						Yes No

## OTHER (Including Author, Title, Date, Pertinent Pages, etc.)

CSM	AN	11	Brown, Robert D. and Martin, Yvonne C., "The Information Content of 2D and 3D Structural Descriptors Relevant to Ligand-Receptor Binding," <i>Journal of Chemical Information and Computer Sciences</i> , American Chemical Society, 1997, Vol. 37, No. 1, pp. 1-9.
	AO	11	Brown, Robert D. and Martin, Yvonne C., "Use of Structure-Activity Data To Compare Structure-Based Clustering Methods and Descriptors for Use in Compound Selection," <i>Journal of Chemical Information and Computer Sciences</i> , American Chemical Society, 1996, Vol. 36, No. 3, pp. 572-584.
	AP	11	Cummins, David J. et al., "Molecular Diversity in Chemical Databases: Comparison of Medicinal Chemistry Knowledge Bases and Databases of Commercially Available Compounds," <i>Journal of Chemical Information and Computer Sciences</i> , American Chemical Society, 1996, Vol. 36, No. 4, pp. 750-763.
	AQ	11	Domine, D. et al., "Non-Linear Mapping for Structure-Activity and Structure-Property Modelling," <i>Journal of Chemometrics</i> , John Wiley & Sons, Ltd., Vol. 7, No. 4, July-August 1993, pp. 227-242.
↓	AR	11	Saunders, M., "Stochastic Exploration of Molecular Mechanics Energy Surfaces. Hunting for the Global Minimum," <i>Journal of the American Chemical Society</i> , American Chemical Society, Vol. 109, 10, May 13, 1987, pages 3150-3152.

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March 16, 2004

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FOURTH SUPPLEMENTAL INFORMATION  
DISCLOSURE STATEMENTATTY. DOCKET NO.  
1503.1070003APPLICATION NO.  
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EXAMINER INITIAL		DOCUMENT NUMBER	DATE	NAME	CLASS	SUB- CLASS	FILING DATE
	AA12						
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EXAMINER INITIAL		DOCUMENT NUMBER	DATE	COUNTRY	CLASS	SUB- CLASS	TRANSLATION
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## OTHER (Including Author, Title, Date, Pertinent Pages, etc.)

CSM	AN	12	Downs, Geoff M. and Barnard, John M., "Techniques for Generating Descriptive Fingerprints in Combinatorial Libraries," <i>Journal of Chemical Information and Computer Sciences</i> , American Chemical Society, 1997, Vol. 37, No. 1, pp. 59-61.
	AO	12	Gillet, Valerie J., "Background Theory of Molecular Diversity," <i>Molecular Diversity in Drug Design</i> , Kluwer Academic Publishers, 1999, pp. 43-65.
	AP	12	Good, Andrew C. and Lewis, Richard A., "New Methodology for Profiling Combinatorial Libraries and Screening Sets: Cleaning Up the Design Process with HARPick," <i>Journal of Medicinal Chemistry</i> , American Chemical Society, 1997, Vol. 40, No. 24, pp. 3926-3936.
	AQ	12	Pal, N.R. and Eluri, V.K., "Two Efficient Connectionist Schemes for Structure Preserving Dimensionality Reduction," <i>IEEE Transactions on Neural Networks</i> , IEEE, Vol 9, No. 6, November 1998, pp. 1142-1154.
↓	AR	12	Jamois, Eric A. et al., "Evaluation of Reagent-Based and Product-Based Strategies in the Design of Combinatorial Library Subsets," <i>Journal of Chemical Information and Computer Sciences</i> , American Chemical Society, 2000, Vol. 40, No.1, pp. 63-70.

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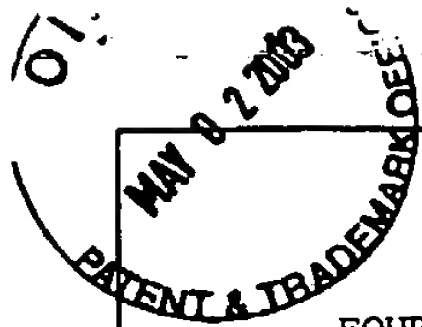
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CSM	AN	13	Kim, H. et al., "Self-Organized Distributed Networks for Learning Highly Nonlinear Mapping," <i>Intelligent Engineering Systems Through Artificial Neural Networks</i> , American Society of Mechanical Engineers, Vol. 4, November 13-16, 1994, pp. 109-114.
	AO	13	Leach, Andrew R. et al., "Implementation of a System for Reagent Selection and Library Enumeration, Profiling, and Design," <i>Journal of Chemical Information and Computer Sciences</i> , American Chemical Society, 1999, Vol. 39, No. 6, pp. 1161-1172.
	AP	13	Lobanov, Victor S. and Agrafiotis, Dimitris K., "Stochastic Similarity Selections from Large Combinatorial Libraries," <i>Journal of Chemical Information and Computer Sciences</i> , American Chemical Society, March/April 2000, Vol. 40, No. 2, pp. 460-470.
	AQ	13	Matter, Hans and Pötter, Thorsten, "Comparing 3D Pharmacophore Triplets and 2D Fingerprints for Selecting Diverse Compound Subsets," <i>Journal of Chemical Information and Computer Sciences</i> , American Chemical Society, 1999, Vol. 39, No. 6, pp. 1211-1225.
↓	AR	13	Matter, Hans, "Selecting Optimally Diverse Compounds from Structure Databases: A Validation Study of Two-Dimensional and Three-Dimensional Molecular Descriptors," <i>Journal of Medicinal Chemistry</i> , American Chemical Society, 1997, Vol. 40, No. 8, pp. 1219-1229.

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FORM PTO-1449 FOURTH SUPPLEMENTAL INFORMATION DISCLOSURE STATEMENT	ATTY. DOCKET NO. 1503.1070003	APPLICATION NO. 09/934,084
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CSM	AN	14	Sadowski, Jens and Kubinyi, Hugo, "A Scoring Scheme for Discriminating between Drugs and Nondrugs," <i>Journal of Medicinal Chemistry</i> , American Chemical Society, 1998, Vol. 41, No. 18, pp. 3325-3329.
	AO	14	Schnur, Dora, "Design and Diversity Analysis of Large Combinatorial Libraries Using Cell-Based Methods," <i>Journal of Chemical Information and Computer Science</i> , American Chemical Society, 1999, Vol. 39, No. 1, pp. 36-45.
	AP	14	Schuffenhauer, Ansgar et al., "Similarity Searching in Files of Three-Dimensional Chemical Structures: Analysis of the BIOSER Database Using Two-Dimensional Fingerprints and Molecular Field Descriptors," <i>Journal of Chemical Information and Computer Science</i> , American Chemical Society, 2000, Vol. 40, No. 2, pp. 295-307.
	AQ	14	Turner, David B. et al., "Rapid Quantification of Molecular Diversity for Selective Database Acquisition," <i>Journal of Chemical Information and Computer Science</i> , American Chemical Society, 1997, Vol. 37, No. 1, pp. 18-22.
✓	AR	14	Wang, Jing and Ramnarayan, Kal, "Toward Designing Drug-Like Libraries: A Novel Computational Approach for Prediction of Drug Feasibility of Compounds," <i>Journal of Combinatorial Chemistry</i> , American Chemical Society, November/December 1999, Vol. 1, No. 6, pp. 524-533.

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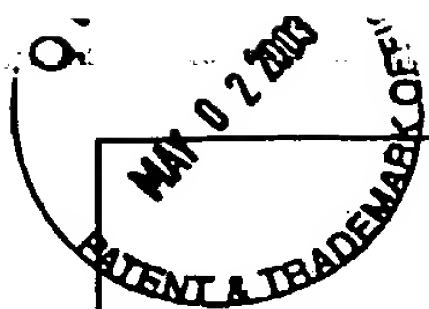
## OTHER (Including Author, Title, Date, Pertinent Pages, etc.)

C9M	AN	15	Gasteiger, J. et al, "Assessment of the Diversity of Combinatorial Libraries by an Encoding of Molecular Surface Properties," <i>Abstracts of Papers Part 1: 211th ACS National Meeting</i> , March 24-28, 1996, p. 70-CINF.
	AO	15	Hassan, Moises et al., "Optimization and visualization of molecular diversity of combinatorial libraries," <i>Molecular Diversity</i> , ESCOM Science Publishers B.V., 1996, Vol. 2, pp. 64-74.
	AP	15	Bellman, R.E., <i>Adaptive Control Processes: A Guided Tour</i> , Princeton Univ. Press, Princeton, NJ (1961), entire book submitted.
	AQ	15	Bezdek, J.C., <i>Pattern Recognition with Fuzzy Objective Function Algorithms</i> , Plenum Press, New York, NY (1981), entire book submitted.
✓	AR	15	Johnson, M.A., and Maggiora, G.M., <i>Concepts and Applications of Molecular Similarity</i> , John Wiley and Sons, New York, NY (1990), entire book submitted.

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	AJ16						No
	AK16						No
	AL16						No
	AM16						No

**OTHER (Including Author, Title, Date, Pertinent Pages, etc.)**

<i>GSM</i>	AN	<u>16</u>	Kohonen, T., <i>Self-Organizing Maps</i> , Springer-Verlag, Heidelberg, Germany (1995), entire book submitted.
	AO	<u>16</u>	Oja, E., <i>Subspace Methods of Pattern Recognition</i> , Research Studies Press Ltd., Letchworth, England (1983), entire book submitted.
	AP	<u>16</u>	Agrafiotis, D.K., "A New Method For Analyzing Protein Sequence Relationships Based On Sammon Maps," <i>Protein Science</i> , Cambridge University Press, Vol. 6, No. 2, February 1997, pp. 287-293.
	AQ	<u>16</u>	Porto, V. et al., "Alternative Neural Network Training Methods," <i>IEEE Expert</i> , IEEE, Vol. 10, No. 4, pages 16-22.
✓	AR	<u>16</u>	Amzel, L.M., "Structure-based drug design," <i>Current Opinion in Biotechnology</i> , Vol. 9, No. 4, August 1998, pp. 366-369.

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	AJ17						No
	AK17						No
	AL17						No
	AM17						No

## OTHER (Including Author, Title, Date, Pertinent Pages, etc.)

CSM	AN	17	Blaney, J.M. and Martin, E.J., "Computational approaches for combinatorial library design and molecular diversity analysis," <i>Current Opinion in Chemical Biology</i> , Current Biology Ltd., Vol. 1, No. 1, June 1997, pp. 54-59.
	AO	17	Mumenthaler, Ch. And Braun, W., "Automated Assignment of Simulated and Experimental NOESY Spectra of Proteins by Feedback Filtering and Self-correcting Distance Geometry," <i>Journal of Molecular Biology</i> , Academic Press Limited, Vol. 254, No. 3, December 1, 1995, pages 465-480.
	AP	17	Cafilisch, A. and Karplus, M., "Computational combinatorial chemistry for de novo ligand design: Review and assessment," <i>Perspectives in Drug Discovery and Design</i> , ESCOM Science Publishers B.V., Vol. 3, 1995, pp. 51-84.
	AQ	17	Meng, E. et al., "Orientational Sampling and Rigid-Body Minimization in Molecular Docking," <i>PROTEINS: Structure, Function and Genetics</i> , Wiley-Liss, Inc., Vol. 17, No. 3, 1993, pages 266-278.
↓	AR	17	Eichler, U. et al., "Addressing the problem of molecular diversity," <i>Drugs of the Future</i> , Prous Science, Vol. 24, No. 2, 1999, pp. 177-190.

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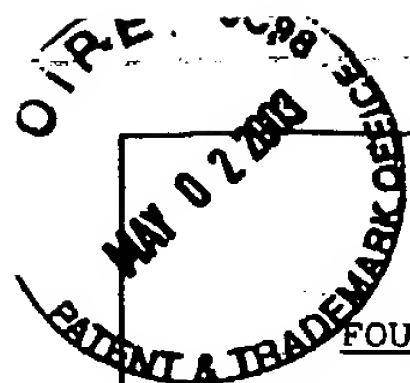
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GSM	AN	18	Felder, E.R. and Poppinger, D., "Combinatorial Compound Libraries for Enhanced Drug Discovery Approaches," <i>Advances in Drug Research</i> , Academic Press, Vol. 30, 1997, pp. 112-199.
	AO	18	Geysen, H.M. and Mason, T.J., "Screening Chemically Synthesized Peptide Libraries for Biologically-Relevant Molecules," <i>Bioorganic &amp; Medicinal Chemistry Letters</i> , Pergamon Press Ltd., Vol. 3, No. 3, 1993, pp. 397-404.
	AP	18	Gobbi, A. et al., "New Leads By Selective Screening of Compounds From Large Databases," <i>Abstracts of Papers Part 1: 213th ACS National Meeting</i> , American Chemical Society, April 13-17, 1997, p. 67-CINF.
	AQ	18	Houghten, R.A. et al., "The Use of Synthetic Peptide Combinatorial Libraries for the Identification of Bioactive Peptides," <i>Peptide Research</i> , Vol. 5, No. 6, 1992, pp. 351-358.
✓	AR	18	Klopman, G., "Artificial Intelligence Approach to Structure-Activity Studies. Computer Automated Structure Evaluation of Biological Activity of Organic Molecules," <i>Journal of the American Chemical Society</i> , American Chemical Society, Vol. 106, No. 24, 1984, pp. 7315-7321.

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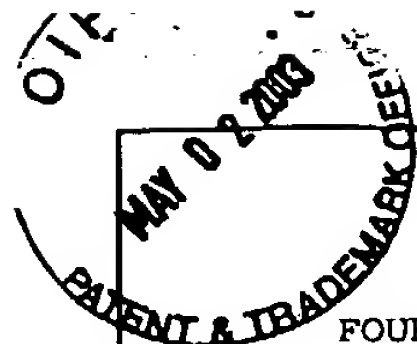
CSM	AN	19	Lajiness, M.S. et al., "Implementing Drug Screening Programs Using Molecular Similarity Methods," QSAR: Quantitative Structure-Activity Relationships in Drug Design, Alan R. Liss, Inc., 1989, pp. 173-176.
	AO	19	Loew, G.H. et al., "Strategies for Indirect Computer-Aided Drug Design," <i>Pharmaceutical Research</i> , Plenum Publishing Corporation, Vol. 10, No. 4, 1993, pp. 475-486.
	AP	19	Lynch, M.F. et al., "Generic Structure Storage and Retrieval," <i>Journal of Chemical Information and Computer Sciences</i> , American Chemical Society, Vol. 25, No. 3, August 1985, pp. 264-270.
	AQ	19	Myers, P.L. et al., "Rapid, Reliable Drug Discovery," <i>Today's Chemist At Work</i> , American Chemical Society, Vol. 6, No. 7, July/August 1997, pp. 46-48, 51 & 53.
✓	AR	19	Pabo, C.O. and Suchanek, E.G., "Computer-Aided Model-Building Strategies for Protein Design," <i>Biochemistry</i> , American Chemical Society, Vol. 25, No. 20, 1986, pp. 5987-5991.

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	AJ20						Yes No
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CSM	AN	20	Saudek, V. et al., "Solution Conformation of Endothelin-1 by H NMR, CD, and Molecular Modeling," <i>International Journal of Peptide Protein Research</i> , Munksgaard International Publishers Ltd., Vol. 37, No. 3, 1991, pp. 174-179.
	AO	20	Singh, J. et al., "Application of Genetic Algorithms to Combinatorial Synthesis: A Computational Approach to Lead Identification and Lead Optimization," <i>J. Am. Chem. Soc.</i> , American Chemical Society, Vol. 118, No. 7, February 7, 1996, pp. 1669-1676.
	AP	20	Van Drie, J.H. and Lajiness, M.S., "Approaches to virtual library design," <i>Drug Discovery today</i> , Elsevier Science Ltd., Vol. 3, No. 6, June 1998, pp. 274-283.
	AQ	20	Walters, W.P. et al., "Virtual screening - an overview," <i>Drug Discovery today</i> , Elsevier Science Ltd., Vol. 3, No. 4, April 1998, pp. 160-178.
✓	AR	20	Weber, L., "Evolutionary combinatorial chemistry: application of genetic algorithms," <i>Drug Discovery today</i> , Elsevier Science Ltd., Vol. 3, No. 8, August 1998, pp. 379-385.

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	AK21						Yes No
	AL21						Yes No
	AM21						Yes No

## OTHER (Including Author, Title, Date, Pertinent Pages, etc.)

CSM	AN	21	Weber, L. et al., "Optimization of the Biological Activity of Combinatorial Compound Libraries by a Genetic Algorithm," <i>Angewandte Chemie International Edition in English</i> , VCH, Vol. 34, No. 20, November 3, 1995, pp. 2280-2282.
	AO	21	Graybill, T.L. et al., "Enhancing the Drug Discovery Process by Integration of High-Throughput Chemistry and Structure-Based Drug Design," <i>Molecular Diversity and Combinatorial Chemistry: Libraries and Drug Discovery</i> , American Chemical Society, 1996, pp. 16-27.
	AP	21	Saund, E., "Dimensionality-Reduction Using Connectionist Networks," <i>IEEE Transactions on Pattern Analysis and Machine Intelligence</i> , IEEE, Vol. 11, No. 3, March 1989, pp. 304-314.
	AQ	21	"3DP gains drug research patent", <i>Chemistry in Britain</i> , The Royal Society of Chemistry, Vol. 32, No. 1, January 1996, p. 22.
V	AR	21	"Accelerate the Discovery Cycle with Chem-XI", Source and date of publication unclear, 2 pages.

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	AK22						Yes No
	AL22						Yes No
	AM22						Yes No

## OTHER (Including Author, Title, Date, Pertinent Pages, etc.)

CSM	AN	22	Agrafiotis, D. K., "Stochastic Algorithms for Maximizing Molecular Diversity", <i>Journal of Chemical Information and Computer Sciences</i> , American Chemical Society, Vol. 37, No. 5, 1997, pp. 841-851.
	AO	22	Alsberg, B.K. et al., "Classification of pyrolysis mass spectra by fuzzy multivariate rule induction-comparison with regression, K-nearest neighbour, neural and decision-tree methods", <i>Analytica Chimica Acta</i> , Elsevier Science B.V., Vol. 348, No. 1-3, August 20, 1997, pp. 389-407.
	AP	22	Andrea, T.A. and Kalayeh, H., "Applications of Neural Networks in Quantitative Structure-Activity Relationships of Dihydrofolate Reductase Inhibitors", <i>Journal of Medicinal Chemistry</i> , American Chemical Society, Vol. 34, No. 9, 1991, pp. 2824-2836.
	AQ	22	Aoyama, T. et al., "Neural Networks Applied to Quantitative Structure-Activity Relationship Analysis", <i>Journal of Medicinal Chemistry</i> , American Chemical Society, Vol. 33, No. 9, 1990, pp. 2583-2590.
	AR	22	Aoyama, T. and Ichikawa, H., "Obtaining the Correlation Indices between Drug Activity and Structural Parameters Using a Neural Network", <i>Chemical &amp; Pharmaceutical Bulletin</i> , Pharmaceutical Society of Japan, Vol. 39, No. 2, February 1991, pp. 372-378.

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CSM	AN	23	Leach, A., "A Survey of Methods for Searching the Conformational Space of Small and Medium-Sized Molecules," <i>Reviews in Computational Chemistry</i> , VCH Publishers, Vol. 2, pages 1-55.
	AO	23	Baum, R.M., "Combinatorial Approaches Provide Fresh Leads for Medicinal Chemistry", <i>Chemical &amp; Engineering News</i> , American Chemical Society, February 7, 1994, pp. 20-26.
	AP	23	Bentley, J. L., "Multidimensional Binary Search Trees Used for Associative Searching", <i>Communications of the ACM</i> , Association for Computing Machinery, Inc., Vol. 18, No. 9, September 1975, pp. 509-517.
	AQ	23	Bottou, L. and Vapnik, V. "Local Learning Algorithms", <i>Neural Computation</i> , Massachusetts Institute of Technology, Vol. 4, No. 6, November 1992, pp. 888-900.
	AR	23	Boulu, L.G. and Crippen, G.M., "Voronoi Binding Site Models: Calculation of Binding Modes and Influence of Drug Binding Data Accuracy", <i>Journal of Computational Chemistry</i> , John Wiley & Sons, Inc., Vol. 10, No. 5, July/August 1989, pp. 673-682.

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March 16, 2004

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CSM	AN	24	Boulu, L.G. et al., "Voronoi Binding Site Model of a Polycyclic Aromatic Hydrocarbon Binding Protein", <i>Journal of Medicinal Chemistry</i> , American Chemical Society, Vol. 33, No. 2, 1990, pp. 771-775.
	AO	24	Cacoullos, T., "Estimation of a Multivariate Density", <i>Annals of The Institute of Statistical Mathematics</i> , The Institute of Statistical Mathematics, Vol. 18, No. 2, 1966, pp. 179-189.
	AP	24	Clark, R.D., "OptiSim: An Extended Dissimilarity Selection Method for Finding Diverse Representative Subsets", <i>Journal of Chemical Information and Computer Sciences</i> , American Chemical Society, Vol. 37, No. 6, 1997, pp. 1181-1188.
	AQ	24	Clark, D. E., and Westhead, D.R., "Evolutionary algorithms in computer-aided molecular design", <i>Journal of Computer-Aided Molecular Design</i> , ESCOM Science Publishers B.V., Vol. 10, No. 4, August 1996, pp. 337-358.
✓	AR	24	Cramer, III, R. D. et al., "Comparative Molecular Field Analysis (CoMFA). 1. Effect of Shape on Binding of Steroids to Carrier Proteins", <i>Journal of The American Chemical Society</i> , American Chemical Society, Vol. 110, No. 18, August 31, 1988, pp. 5959-5967.

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	AK25						Yes No
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CSM	AN	25	Cramer, III, R. D. et al., "Substructural Analysis. A Novel Approach to the Problem of Drug Design", <i>Journal of Medicinal Chemistry</i> , Vol. 17, No. 5, May 1974, pp. 533-535.
	AO	25	Crippen, G. M., "Voronoi Binding Site Models", <i>Journal of Computational Chemistry</i> , John Wiley & Sons, Inc., Vol. 8, No. 7, October/November 1987, pp. 943-955.
	AP	25	Friedman, J. H. et al., "An Algorithm for Finding Best Matches in Logarithmic Expected Time", <i>ACM Transactions on Mathematical Software</i> , Association for Computing Machinery, Vol. 3, No. 3, September 1977, pp. 209-226.
	AQ	25	Friedman, J.H., "Fitting Functions To Noisy Data In High Dimensions", Department of Statistics- Stanford University Technical Report No. 101, (August, 1988), pages 1-36.
↓	AR	25	Gallop, M. A. et al., "Applications of Combinatorial Technologies to Drug Discovery. 1. Background and Peptide Combinatorial Libraries", <i>Journal of Medicinal Chemistry</i> , American Chemical Society, Vol. 37, No. 9, April 29, 1994, pp. 1233-1251.

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	AJ26						Yes No
	AK26						Yes No
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## OTHER (Including Author, Title, Date, Pertinent Pages, etc.)

CSM	AN	26	Ghose, A. K. and Crippen, G.M., "Use of Physicochemical Parameters in Distance Geometry and Related Three-Dimensional Quantitative Structure-Activity Relationships: A Demonstration Using <i>Escherichia coli</i> Dihydrofolate Reductase Inhibitors", <i>Journal of Medicinal Chemistry</i> , American Chemical Society, Vol. 28, No. 3, 1985, pp. 333-346.
	AO	26	Good, A. C. et al., "Structure-Activity Relationships from Molecular Similarity Matrices", <i>Journal of Medicinal Chemistry</i> , American Chemical Society, Vol. 36, No. 4, February 19, 1993, pp. 433-438.
	AP	26	Gordon, E. M. et al., "Applications of Combinatorial Technologies to Drug Discovery. 2. Combinatorial Organic Synthesis, Library Screening Strategies, and Future Directions", <i>Journal of Medicinal Chemistry</i> , American Chemical Society, Vol. 37, No. 10, May 13, 1994, pp. 1385-1401.
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↓	AR	26	Hopfinger, A. J., "A QSAR Investigation of Dihydrofolate Reductase Inhibition by Baker Triazines Based upon Molecular Shape Analysis", <i>Journal of the American Chemical Society</i> , American Chemical Society, Vol. 102, No. 24, November 19, 1980, pp. 7196-7206.

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	AJ27						Yes No
	AK27						Yes No
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	AM27						Yes No

## OTHER (Including Author, Title, Date, Pertinent Pages, etc.)

CA01	AN	27	Jackson, R. C., "Update on computer-aided drug design", <i>Current Opinion in BIOTECHNOLOGY</i> , Current Biology Ltd., Vol. 6, No. 6, December 1995, pp. 646-651.
	AO	27	Kim, K. H., "Comparative molecular field analysis (CoMFA)", <i>Molecular Similarity in Drug Design</i> , ed. P. M. Dean, Blackie Academic & Professional, 1995, Ch. 12, pp. 291-331.
	AP	27	Kohonen, T., "Self-Organized Formation of Topologically Correct Feature Maps", <i>Biological Cybernetics</i> , Springer-Verlag, Vol. 43, No. 1, 1982, pp. 59-69.
	AQ	27	Koile, K. and Shapiro, R., "Building A Collaborative Drug Design System", <i>Proceedings of the 25th Hawaii International Conference on System Sciences</i> , IEEE, 1992, pp. 706-716.
↓	AR	27	Kowalski, B. R. and Bender, C. F., "Pattern Recognition. II. Linear and Nonlinear Methods for Displaying Chemical Data", <i>Journal of the American Chemical Society</i> , American Chemical Society, Vol. 95, No. 3, February 7, 1973, pp. 686-693.

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	AJ28						Yes No
	AK28						Yes No
	AL28						Yes No
	AM28						Yes No

## OTHER (Including Author, Title, Date, Pertinent Pages, etc.)

CSM	AN	<u>28</u>	Kruskal, J. B., "Nonmetric Multidimensional Scaling: A Numerical Method", <i>Psychometrika</i> , Vol. 29, No. 2, June, 1964, pp. 115-129.
	AO	<u>28</u>	Lengauer, T. and Rarey, M., "Computational methods for biomolecular docking", <i>Current Opinion in Structural Biology</i> , Current Biology Ltd, Vol. 6, No. 3, June, 1996, pp. 402-406.
	AP	<u>28</u>	Luke, B. T., "Evolutionary Programming Applied to the Development of Quantitative Structure-Activity Relationships and Quantitative Structure-Property Relationships", <i>Journal of Chemical Information and Computer Sciences</i> , American Chemical Society, Vol. 34, No. 6, November/December 1994, pp. 1279-1287.
	AQ	<u>28</u>	Martin, E. J. et al., "Does Combinatorial Chemistry Obviate Computer-Aided Drug Design?", <i>Reviews in Computational Chemistry</i> , VCH Publishers, Inc., Vol. 10, 1997, pp. 75-99.
✓	AR	<u>28</u>	Martin, E. J. et al., "Measuring Diversity: Experimental Design of Combinatorial Libraries for Drug Discovery", <i>Journal of Medicinal Chemistry</i> , American Chemical Society, Vol. 38, No. 9, April 28, 1995, pp. 1431-1436.

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	AK29						Yes No
	AL29						Yes No
	AM29						Yes No

## OTHER (Including Author, Title, Date, Pertinent Pages, etc.)

CSM	AN	29	McMartin, C. and Bohacek, R.S., "QXP: Powerful, rapid computer algorithms for structure-based drug design", <i>Journal of Computer-Aided Molecular Design</i> , Kluwer Academic Publishers, Vol. 11, No. 4, July 1997, pp. 333-344.
	AO	29	Mezey, P. G. and Walker, P.D., "Fuzzy molecular fragments in drug research", <i>Drug Discovery today</i> , Vol. 2, No. 4, April 1997, pp. 132-137.
	AP	29	Müller, K., "On the paradigm shift from rational to random design", <i>Journal of Molecular Structure (Theochem)</i> , Elsevier Science B.V., Vol. 398-399, Special Issue, 1997, pp. 467-471.
	AQ	29	Jorgensen, W. and Tirado-Rives, J., "Monte Carlo vs. Molecular Dynamics for Conformational Sampling," <i>Journal of Physical Chemistry</i> , American Chemical Society, Vol. 100, No. 34, August 22, 1996, pages 14508-14513.
↓	AR	29	Kuszewski, J. et al., "Sampling and efficiency of metric matrix, distance geometry: A novel partial metrization algorithm," <i>Journal of Biomolecular NMR</i> , Escom Science Publishers B.V., Vol. 2, No. 1, January 1992, pages 33-56.

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	AM30						Yes No

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CGM	AN	30	Omohundro, S. M., "Bumptrees for Efficient Function, Constraint, and Classification Learning", <i>Advances in Neural Information Processing Systems 3</i> , Morgan Kaufmann, 1991, 7 pages, unknown.
	AO	30	Parrill, A. L., "Evolutionary and genetic methods in drug design", <i>Drug Discovery today</i> , Elsevier Science Ltd., Vol. 1, No. 12, December 1996, pp. 514-521.
	AP	30	Polanski, J., "A neural network for the simulation of biological systems", <i>Journal of Molecular Structure (Theochem)</i> , Elsevier Science Ltd., Vol. 398-399, Special Issue, 1997, pp. 565-571.
	AQ	30	Ramos-Nino, M. E. et al., "A comparison of quantitative structure-activity relationships for the effect of benzoic and cinnamic acids on <i>Listeria monocytogenes</i> using multiple linear regression, artificial neural network and fuzzy systems", <i>Journal of Applied Microbiology</i> , Society for Applied Bacteriology, Vol. 82, No. 2, February 1997, pp. 168-176.
✓	AR	30	Rogers, D. and Hopfinger, A. J., "Application of Genetic Function Approximation to Quantitative Structure-Activity Relationships and Quantitative Structure-Property Relationships", <i>Journal of Chemical Information and Computer Sciences</i> , American Chemical Society, Vol. 34, No. 4, July/August 1994, pp. 854-866.

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COM	AN	31	Sammon, Jr., J. W., "A Nonlinear Mapping for Data Structure Analysis", <i>IEEE Transactions on Computers</i> , IEEE, Vol. C-18, No. 5, May 1969, pp. 401-409.
	AO	31	Simon, Z. et al., "Mapping of Dihydrofolate-reductase Receptor Site by Correlation with Minimal Topological (Steric) Differences", <i>Journal of Theoretical Biology</i> , Academic Press, Inc., Vol. 66, No. 3, June 7, 1997, pp. 485-495.
	AP	31	Smellie, A. S. et al., "Fast Drug-Receptor Mapping by Site-Directed Distances: A Novel Method of Predicting New Pharmacological Leads", <i>Journal of Chemical Information and Computer Sciences</i> , American Chemical Society, Vol. 31, No.3, August 1991, pp. 386-392.
	AQ	31	Specht, D. F., "A General Regression Neural Network", <i>IEEE Transactions on Neural Networks</i> , IEEE, Vol. 2, No. 6, November 1991, pp. 568-576.
✓	AR	31	Svozil, D. et al., "Neural Network Prediction of the Solvatochromic Polarity/Polarizability Parameter $\pi^H_2$ ", <i>Journal of Chemical Information and Computer Sciences</i> , American Chemical Society, Vol. 37, No. 2, 1997, pp. 338-342.

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	AO	32	Torgerson, W. S., "Multidimensional Scaling: I. Theory and Method", <i>Psychometrika</i> , The Psychometric Society, Vol. 17, No. 4, December 1952, pp. 401-419.
	AP	32	Vapnik, V., "Principles of Risk Minimization for Learning Theory", <i>Advances in Neural Information Processing Systems 4</i> , Morgan Kaufmann Publishers, Inc., 1992, pp. 831-838.
	AQ	32	Vapnik, V. and Bottou, L., "Local Algorithms for Pattern Recognition and Dependencies Estimation", <i>Neural Computation</i> , Massachusetts Institute of Technology, Vol. 5, No. 6, November 1993, pp. 893-909.
✓	AR	32	Viswanadhan, V. N. et al., "Mapping the binding site of the nucleoside transporter protein: a 3D-QSAR study", <i>Biochimica et Biophysica Acta</i> , Elsevier Science Publishers B.V., Vol. 1039, No. 3, 1990, pp. 356-366.

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CSM	AN	33	Jain, A. <i>et al.</i> , "Artificial Neural Networks: A Tutorial," IEEE, March 1996, pages 31-44.
	AO	33	Westhead, D. R. <i>et al.</i> , "A comparison of heuristic search algorithms for molecular docking", <i>Journal of Computer-Aided Molecular Design</i> , Kluwer Academic Publishers, Vol. 11, 1997, pp. 209-228.
	AP	33	Willett, P., "Genetic algorithms in molecular recognition and design", <i>Trends in Biotechnology</i> , Elsevier Science Publishers B.V., Vol. 13, No. 12, December 1995, pp. 516-521.
	AQ	33	Willett, P. and Winterman, V., "A Comparison of Some Measures for the Determination of Inter-Molecular Structural Similarity Measures of Inter-Molecular Structural Similarity", <i>Quantitative Structure-Activity Relationships</i> , VCH, Vol. 5, No. 1, March 1986, pp. 18-25.
↓	AR	33	Zadeh, L. A., "Communication Fuzzy Algorithms", <i>Information and Control</i> , Academic Press Inc., Vol. 12, No. 2, February 1968, pp. 94-102.

EXAMINER

DATE CONSIDERED

EXAMINER: Initial if reference considered, whether or not citation is in conformance with MPEP 609. Draw line through citation if not in conformance and not considered. Include copy of this form with next communication to Applicant.

FORM PTO-1449

FOURTH SUPPLEMENTAL INFORMATION  
DISCLOSURE STATEMENTATTY. DOCKET NO.  
1503.1070003APPLICATION NO.  
09/934,084APPLICANTS  
Lobanov et al.FILING DATE  
August 22, 2001GROUP  
1631

## U.S. PATENT DOCUMENTS

EXAMINER INITIAL	DOCUMENT NUMBER	DATE	NAME	CLASS	SUB- CLASS	FILING DATE
	AA34					
	AB34					
	AC34					
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## FOREIGN PATENT DOCUMENTS

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	AK34					Yes No
	AL34					Yes No
	AM34					Yes No

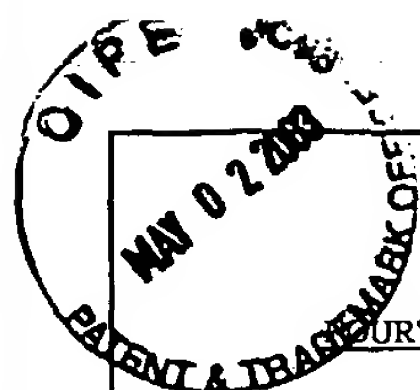
## OTHER (Including Author, Title, Date, Pertinent Pages, etc.)

CSM	AN	34	Zadeh, L. A., "Fuzzy Sets", <i>Information and Control</i> , Academic Press Inc., Vol. 8, No. 3, June 1965, pp. 338-353.
	AO	34	Havel, T., "A New Method for Building Protein Conformations from Sequence Alignments with Homologues of Known Structure," <i>Journal of Molecular Biology</i> , Academic Press Limited, Vol 217, No. 1, January 5, 1991, pages 1-7.
	AP	34	Havel, T. and Wüthrich, K., "A Distance Geometry Program for Determining the Structures of Small Proteins and other Macromolecules from Nuclear Magnetic Resonance Measurements of Intramolecular <sup>1</sup> H- <sup>1</sup> H Proximities in Solution," <i>Bulletin of Mathematical Biology</i> , Pergamon Press, Vol. 46, No. 4, 1984, pages 673-698.
	AQ	34	Aoyama, T. et al., "Neural Networks Applied to Structure-Activity Relationships," <i>Journal of Medicinal Chemistry</i> , American Chemical Society, Vol. 33., No. 3, 1990, pp. 905-908.
✓	AR	34	Gasteiger, J. et al., "Analysis of the Reactivity of Single Bonds in Aliphatic Molecules by Statistical and Pattern Recognition Methods," <i>Journal of Chemical Information and Computer Sciences</i> , American Chemical Society, Vol. 33, No. 3, 1993, pp. 385-394.

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	AK35						Yes No
	AL35						Yes No
	AM35						Yes No

## OTHER (Including Author, Title, Date, Pertinent Pages, etc.)

GM	AN	35	Guez, A. and Nevo, I., "Neural networks and fuzzy logic in clinical laboratory computing with application to integrated monitoring," <i>Clinica Chimica Acta</i> , Elsevier Science Publishers B.V., Vol. 248, 1996, pp. 73-90.
	AO	35	Rouvray, D.H., "Similarity in Chemistry: Past, Present and Future," <i>Topics in Chemistry</i> , Springer-Verlag, Vol. 173, 1995, pp. 1-30.
	AP	35	de Ridder, D. and Duin, R.P.W., "Sammon's mapping using neural networks: A comparison," <i>Pattern Recognition Letters</i> , Elsevier Science Publishers B.V., Vol. 18, No. 11-13, 1997, pp. 1307-1316.
	AQ	35	Havel, T. and Wüthrich, K., "An Evaluation of the Combined Use of Nuclear Magnetic Resonance and Distance Geometry for the Determination of Protein Conformations in Solution," <i>Journal of Molecular Biology</i> , Academic Press Inc., Vol. 182, No. 2, March 20, 1985, pages 281-294.
✓	AR	35	Chang, G. et al., "An Internal Coordinate Monte Carlo Method for Searching Conformational Space," <i>Journal of the American Chemical Society</i> , American Chemical Society, Vol. III, June 1989, No. 12, pages 4379-4386.

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	AJ36						Yes No
	AK36						Yes No
	AL36						Yes No
	AM36						Yes No

## OTHER (Including Author, Title, Date, Pertinent Pages, etc.)

CSM	AN	36	Crippen, G.M. and Havel, T.F., <i>Distance Geometry and Molecular Conformation</i> , Research Studies Press Ltd., 1988, entire book submitted.
	AO	36	Feuston, B. et al., "Comparison of Knowledge-Based and Distance Geometry Approaches for Generation of Molecular Conformations," <i>Journal of Information and Computer Sciences</i> , American Chemical Society, Vol. 41, No. 3, 2001, pages 754-763.
	AP	36	Ferguson, D. and Raber, D., "A New Approach to Probing Conformational Space with Molecular Mechanics: Random Incremental Pulse Search," <i>Journal of the American Chemical Society</i> , American Chemical Society, Vol. 111, No. 12, 1989, pages 4371-4378.
	AQ	36	Halgren, T. and Nachbar, R., "Merck Molecular Force Field. IV. Conformational Energies and Geometries for MMFF94*," <i>Journal of Computational Chemistry</i> , John Wiley & Sons, Inc., Vol. 17, Nos. 5 & 6, 1996, pages 587-915.
✓	AR	36	Halgren, T., "Merck Molecular Force Field. V. Extension of MMFF94 Using Experimental Data, Additional Computational Data, and Empirical Rules*," <i>Journal of Computational Chemistry</i> , John Wiley & Sons, Inc., Vol. 17, Nos. 5 & 6, April 1996, pages 616-641.

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## FOREIGN PATENT DOCUMENTS

EXAMINER INITIAL		DOCUMENT NUMBER	DATE	COUNTRY	CLASS	SUB- CLASS	TRANSLATION
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	AK37						Yes No
	AL37						Yes No
	AM37						Yes No

## OTHER (Including Author, Title, Date, Pertinent Pages, etc.)

CSM	AN	<u>37</u>	Huang, E. et al., "Distance geometry generates native-like folds for small helical proteins using the consensus distances of predicted protein structures," <i>Protein Science</i> , The Protein Society, Vol. 7, No. 9, September 1998, pages 1998-2003.
	AO	<u>37</u>	
	AP	<u>37</u>	
	AQ	<u>37</u>	
	AR	<u>37</u>	

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## ELECTRONIC INFORMATION DISCLOSURE STATEMENT

Electronic Version v18  
Stylesheet Version v18.0

Title of  
Invention

Method, System and Computer Program Product for  
Determining Properties of Combinatorial Library Products  
from Features of Library Building Blocks

Application Number: 09/934084

Confirmation Number: 7373

First Named Applicant: Victor Lobanov

Attorney Docket Number: 1503.1070003

Art Unit: 1631

Search string: ( 5789160 or 5807754 or 5811241 or 5832494  
or 5858660 or 5861532 or 5866334 or 5901069  
or 5908960 or 5933819 or 6014661 or 6026397  
or 6037135 or 6049797 or 6185506 ).pn.

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### US Patent Documents

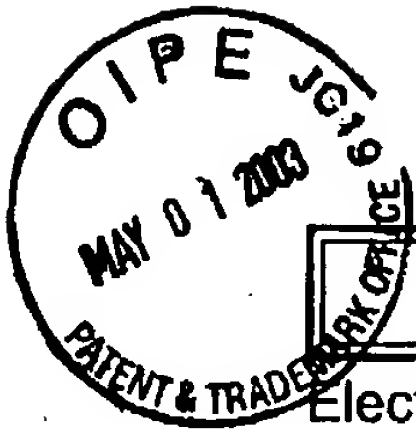
Note: Applicant is not required to submit a paper copy of cited US Patent Documents

init	Cite.No.	Patent No.	Date	Patentee	Kind	Class	Subclass
✓	1	5789160	1998-08-04	Eaton et al.		435	6
	2	5807754	1998-09-15	Zambias et al.		436	518
	3	5811241	1998-09-22	Goodfellow et al.		435	7.1
	4	5832494	1998-11-03	Egger et al.		707	102
	5	5858660	1999-01-12	Eaton et al.		435	6
	6	5861532	1999-01-19	Brown et al.		584	123
	7	5866334	1999-02-02	Beutel		435	6
	8	5901069	1999-05-04	Agrafiotis et al.		384	528.03
	9	5908960	1999-06-01	Newlander		584	177
	10	5933819	1999-08-03	Skolnick et al.		706	21
	11	6014661	2000-01-11	Ahlberg et al.		707	3
	12	6026397	2000-02-15	Sheppard		707	5
	13	6037135	2000-03-14	Kubo et al.		435	7.24
	14	6049797	2000-04-11	Guha et al.		707	6
✓	15	6185506	2001-02-06	Cramer et al.		702	19



Signature

Examiner Name	Date
<i>C. M. 18</i>	<i>March 16, 2004</i>



## ELECTRONIC INFORMATION DISCLOSURE STATEMENT

Electronic Version v18

Stylesheet Version v18.0

Title of  
Invention

Method, System and Computer Program Product for  
Determining Properties of Combinatorial Library Products  
from Features of Library Building Blocks

Application Number: 09/934084

Confirmation Number: 7373

First Named Applicant: Victor Lobanov

Attorney Docket Number: 1503.1070003

Art Unit: 1631

Search string: ( 4773099 or 4811217 or 4859736 or 4908773  
or 4935875 or 4939666 or 5010175 or 5025388  
or 5155801 or 5167009 or 5181259 or 5240680  
or 5260882 or 5265030 or 5270170 or 5288514  
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### US Patent Documents

Note: Applicant is not required to submit a paper copy of cited US Patent Documents

init	Cite.No.	Patent No.	Date	Patentee	Kind	Class	Subclass
✓	1	4773099	1988-09-20	Bokser		382	14
	2	4811217	1989-03-07	Tokizane et al.		364	300
	3	4859736	1989-08-22	Rink		525	54.1
	4	4908773	1990-03-13	Pantoliano et al.		364	496
	5	4935875	1990-06-19	Shah et al.		364	497
	6	4939666	1990-07-03	Hardman		364	496
	7	5010175	1991-04-23	Rutter et al.		530	334
✓	8	5025388	1991-06-18	Cramer, III et al.		364	496

✓	9	5155801	1992-10-13	Lincoln	395	22
	10	5167009	1992-11-24	Skeirik	395	27
	11	5181259	1993-01-19	Rorvig	382	36
	12	5240680	1993-08-31	Zuckermann et al.	422	67
	13	5260882	1993-11-09	Blanco et al.	364	499
	14	5265030	1993-11-23	Skolnick et al.	364	496
	15	5270170	1993-12-14	Schatz et al.	435	7.37
	16	5288514	1994-02-22	Ellman	427	2
	17	5307287	1994-04-26	Cramer, III et al.	364	496
	18	5323471	1994-06-21	Hayashi	382	15
	19	5331573	1994-07-19	Balaji et al.	364	500
	20	5434796	1995-07-18	Weininger	364	496
	21	5436850	1995-07-25	Eisenberg et al.	364	496
	22	5442122	1995-08-15	Noda et al.	564	426
	23	5463564	1995-10-31	Agrafiotis et al.	364	496
	24	5499193	1996-03-12	Sugawara et al.	364	500
	25	5519635	1996-05-21	Miyake et al.	364	497
	26	5524065	1996-06-04	Yagasaki	382	226
	27	5526281	1996-06-11	Chapman et al.	364	496
	28	5545568	1996-08-13	Ellman	436	518
	29	5549974	1996-08-27	Holmes	428	403
	30	5095443	1992-03-10	Watanabe	364	513
	31	5553225	1996-09-03	Perry	395	157
	32	5565325	1996-10-15	Blake	435	7.1
	33	5574656	1996-11-12	Agrafiotis et al.	364	500
	34	5585277	1996-12-17	Bowie et al.	436	518
	35	5598510	1997-01-28	Castelaz	395	23
	36	5602755	1997-02-11	Ashe et al.	364	498
	37	5602938	1997-02-11	Akiyama et al.	382	155
	38	5612895	1997-03-18	Balaji et al.	364	496
	39	5621861	1997-04-15	Hayashi et al.	395	23
	40	5634017	1997-05-27	Mohanty et al.	395	326
	41	5635598	1997-06-03	Lebl et al.	530	334
	42	5670326	1997-09-23	Beutel	435	7.1
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Signature

Examiner Name	Date
<i>C. M. A.</i>	<i>March 16, 2004</i>



FORM PTO-1449

## INFORMATION DISCLOSURE STATEMENT

ATTY. DOCKET NO.  
1503.1070003APPLICATION NO.  
09/934,084APPLICANTS  
Lobanov et al.FILING DATE  
August 22, 2001GROUP  
1631

## U.S. PATENT DOCUMENTS

EXAMINER INITIAL		DOCUMENT NUMBER	DATE	NAME	CLASS	SUB- CLASS	FILING DATE
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## FOREIGN PATENT DOCUMENTS

EXAMINER INITIAL		DOCUMENT NUMBER	DATE	COUNTRY	CLASS	SUB- CLASS	TRANSLATION
CSM	AL1	WO 98/20437, A2	05/1998	PCT	606F	17/50	Yes No
	AM1	WO 94/28504 / A1	12/1994	PCT	606F	15/80	Yes No
↓	AN1	WO 99/35599 / A1	07/1999	PCT	606F	17/50	Yes No
	AO1						Yes No
	AP1						Yes No

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	AS	<u>1</u>	
	AT	<u>1</u>	

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ATTY. DOCKET NO.  
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APPLICATION NO.  
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APPLICANTS  
Lobanov et al.

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U.S. PATENT DOCUMENTS

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FOREIGN PATENT DOCUMENTS

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	AM1						Yes No
	AN1						Yes No
	AO1						Yes No
	AP1						Yes No

OTHER (Including Author, Title, Date, Pertinent Pages, etc.)

CSM	AR	1	Linusson et al., "Statistical Molecular Design of Building Blocks for Combinatorial Chemistry," Journal of Medical Chemistry, Vol. 43, No. 7, American Chemical Society, Published on Web 03/08/2000, pp. 1320-1328.
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	AT	1	

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*C. M. 17*

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*March 16, 2004*

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